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# **EUROPEAN PATENT APPLICATION**

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- Organoclay compositions manufactured with organic acid ester-derived quaternary ammonium compounds, their preparation and non-aqueous fluid systems containing such compositions
- (57) An organophilic clay comprising the reaction product of
  - (a) a smectite-type clay; and
  - (b) a quaternary ammonium compound or compounds selected from the group consisting of:



wherein R, is an alkyl or aralkyl- ester group having 8 to 30 carbon atoms and Ro, Ro and Ra are independently selected from the group consisting of (i) R<sub>1</sub>, (ii) long chain linear or branched alkyl (including methyl), aliphatic or aromatic groups having 1 to 30 carbon atoms (such groups can also include hydroxylated groups); (iii) aralkyl groups, which are benzyl and substituted benzyl moieties, including such groups having fused ring moieties having linear chains or branches of 1 to 30 carbon atoms; (iv) aryl groups such as phenyl and substituted phenyl including fused ring aromatic substituents; (v) beta. gamma unsaturated groups having six or less carbon atoms or hydroxyalkyl groups having 2 to 6 carbon atoms; and (vi) hydrogen and M' is an anion selected from the group consisting of chloride. methylsulfate, acetate, iodide and bromide

The organophilic clay is used in a non-aqueous fluid system such as paints, inks, and coatings to provide rheological properties.

Description

## BACKGROUND OF THE INVENTION

## 1. Brief Description of the Field of the Invention:

The present invention relates to novel organophilic clay compositions (hereafter 'organoclay' or 'organophilic clays') which are dispersible in organic or solvent-based i.e. non-aqueous, fluids to provide a wide variety of meological and viscosity-modifier properties to such fluids. These fluids include oil-based parties and coatings as well as oil-based inks, drilling fluids, caulks and adhesives. The invention also pertains to a process for preparing these novel organophilic clays. The invention also includes organic fluid compositions including such organophilic clays are theological additives.

#### 2 Description of the Prior Art.

It has been known since shortly after the Second World War that organophilic clays are useful to thicken a variety of organic liquid compositions. Organophilic clays are modified smectiter-lybe clays prepared by the fraction of an organic cation, usually a qualetrary armmonium chioride compound produced from a fatty nitrile, with a smecifiet-yec clay utilizing various processes known in the aft. Smectite clays, while dispersible in water are not dispersible in organic fluids. If the organic cation contains at least one faity acid group containing 10 or more, preferably 12 or more, carbon atoms, such reaction-product organically processes the ability of dispersing riic, and increasing the viscosity of, organic liquids. Organically, over the last fitty years, have found a large market providing thickening or rheological proporties to an increasingly wide variety of such liquids including paints, costings, inks, adverses and smiller products.

Representative U.S. Patent No. 4,664,820, issued to the assignee hereof, describes the preparation of organophilic clays, acmee of which have become commercial products, that are used to thicken organic compositions. It is also well known that such organically smay function to thicken both polar or non-polar solvents, depending on the substitutions on the organic cation. For purposes of this patent, organic and solvent are used to mean essentially the same thing. Dr. J.W. Jordan, a former senior scientist employed by the assignee hereof, in "Proceedings of the 10th National Conference on Clays and Clay Minerals" (1963), discusses a wide range of applications of organically from high polarity organic industs to low polarity solvent liquids.

More recently, organophilic clay gellants have been developed which are the reaction products of smexities type clays with certain organic cations or mittures of organic cations, and organic anions or anion commissions. These organicality have the advantage of being easily dispersible in particular types of organic and solvent compositions without the need for a dispersion aids or polar activators under normal factory dispersion conditions. Illustrative patients which describe such improved organophilic clays are U. S. Patient Nos. 4, 105.578, 4,208.218, 4.412.018 (issued to one of the conveniency of the instant invention. 4. 450.095 and 4.517.

Recent U.S. patents issued to assignee hereof show various uses of organoclays and processing improvements in making such organoclays using conventional nitrile quartemary ammonium compounds. These patents include U.S. Patent Nos. 4.695.402. 4.829.644; 5.034.105, 5.075.033, and 5.151.155. See also U.S. Patent No. 5.306.647 and U.S. Patent No. 5.306.647 and See also U

Quaternary ammonium compounds produced from fatty ntiriles have their major use as fabric softeners and in industry in sundry operations. In many European countries, the fact that these uses have core under environmental attack has indirectly resulted in decreasing capacity and limited available supplies. The manufacturers of intrife-based quaternary compounds have chasen to reduce capacity by closing selected paints, rather harm until all paints at reduced capacity utilization rates. This roduction in availability is expected to result in an increase in the price of intrife-based quaternary compounds where two five of working and less available.

The disadvantages of most existing commercial organoids compositions for non-aqueous systems include (a) they require the use of minic- processed quaternary ammonium chiorize compounds, which are increasingly becoming inshort supply, and to justify agranoidly compositions lack the biologradable characteristic which is desired by more and more custom as Dedocradablity of the constituents of drilling fluids, particularly those used in ocean oil drilling, including organociays consumed in such fluids, is becoming a requirement of purchasers of such fluids. Organoidays are an important and occannal ingredient in almost all drilling fluids besed on oil chemistry Beyond drilling fluids, almost all organoidays used in contact with the environment, such as those used in cosmotics or household products, increasingly must be blootgradable to yather law or customer desire.

## SUMMARY OF THE INVENTION

A new type of organophilic clay has been discovered in which the guaternary ammonium compound used to make

the organically is made in part from organic acid-derived eaters. These organically show in a number of circumstance either increased efficiency or substantially equivalent performance versus conventional prior and organically and they can be made with quaternary compounds of either present, or anticipated, lower cost. These novel organically shave a different structure with increased X-ray diffraction pattern 600 spacings. The manufacture of these organically includes eliminate the need to use intrile qualetments of the type which European environmental authorities have found to be environmentally unfriendly. The present invention provides an improved, more biodegradable organically large for the standard provides an improved, more biodegradable organically and solvent-based compositions.

Thus, according to one aspect of the invention, an organophilic clay is provided which comprises the reaction product of:

- (a) a smectite-type clay having a cation exchange capacity of at least about 75 milliequivalents per 100 grams of clay, 100% active clay basis;
- (b) one or more organic cations in an amount sufficient to satisfy at least about 75% of the cation exchange capacity of the smedifie-type clay, and of any optional organic anion(s), wherein the cation or cations are quaternary ammonium compound(s) derived from organic acid-derived esters, and optionally
- (c) one or more organic anion(s) that are capable of reacting with the organic cation(s) to form an organic cation/ organic anion ion pair complex which is intercalated with the clay.

The present invention also contemplates a process for preparing an organophilic clay which comprises:

- (a) preparing an aqueous slurry of a smectite-type clay having a cation exchange capacity of at least about 75 milliequivalents per 100 grams of clay, 100% active clay basis;
- (b) heating the slurry to a temperature of from about 20°C to 100°C;
- (c) adding to the slurry:

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- (i) one or more organic cation(s) which are quaternary ammonium compounds derived from organic acid derived esters, and optionally
- (ii) one or more organic anion(s) that are capable of reacting with the organic cation(s) to form an organic cation/organic anion pair which is intercalated with the clay;
- (d) reacting the resulting mixture for a sufficient time to form an organophilic clay; and

(e) recovering the organophilic clay.

The invention also provides non-aqueous organic compositions thickened with the above-indicated organophilic clay. A third aspect of the invention therefore relates to a non-aqueous fluid system which comprises:

- (a) a non-aqueous composition; and
- (b) an organophilic clay comprising the reaction product of:
  - (i) a smectite-type clay having a cation exchange capacity of at least about 75 milliequivalents per 100 grams of clay, 100% active clay basis;
  - (ii) one or more organic cation(s) in an amount sufficient to satisfy at least about 75% of the cation exchange capacity of the smacifie-type clay and of any optional organic anion(s), which are quaternary ammonium compounds derived from organic acid derived esters, and
  - optionally

    (iii) one or more organic anions that are capable of reacting with the clay and/or the organic cation(s) to form

# the above described complex. Description of the Preferred Embodiments:

As stated above, one aspect of the present invention relates to improved, more efficient organophilic clays. These organophilic clays are prepared by reacting a smectite-type clay with an organic cation or cations provided by specific quaternary ammonium compounds.

The clays used to prepare the organophilic clay of this invention are cation-exchangeable smectite clays which have a cation exchange capacity of about 75 or greater milliequivalents per 100 grams of clay, 100 percent active basis (i.e. beneficiated and essentially free of non-clay impurities). Smectite-type clays are well known in science, geology and in the art of rheological additives, and are commercially available from a variety of sources both in the United States and throughout the world. They are unique among clays in that they exhibit the phenomena of swelling to many.

times their size when contacted with water.

Chemical formula descriptions of such natural smectite clays useful in accordance with the present invention are as follows:

## Montmorillonite

where 0.55 ≤ x ≤ 1.10, f ≤ 4 and R is selected from the group consisting of Na, Li, NH₄, and mixtures thereof,

#### Hectorite

where  $0.57 \le x \le 1.15$ ,  $f \le 4$  and R is selected from the group consisting of Na, Li,  $NH_4$ , and mixtures thereof;

#### Bentonite

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where 0 < x < 1.10, 0 < y < 1.10,  $0.55 \le (x + y) \le 1.10$ ,  $1 \le 4$  and R is selected from the group consisting of Na, Li, NH<sub>4</sub> and mixtures thereof:

# Beidellite

$$[AI_{4+v}(Si_{8-x-v}AI_{x+v})0_{20}(OH)_{4-f}F_f]xR+$$

where 0.55 ≤ x ≤ 1.10, 0 ≤ y 0.44, f ≤ 4 and R is selected from the group consisting of Na, Li, NH<sub>4</sub> and mixtures thereof;

#### Stevensite

where 0.28 ≤ x ≤ 0.57, 1 = 4 and R is selected from the group consisting of Na, Li, NH<sub>4</sub>, and mixtures thereof; and

#### Saponite

$$[Mg_{6-y}Al_ySi_{8-x-y}Al_{x+y}O_{20}(OH)_{4-t}F_t]xR^+$$

where 0.58 ≤ x ≤ 1.18, 0 ≤ y ≤ 0.66, 1 ≤ 4 and R is selected from the group consisting of Na, Li, NH<sub>a</sub>, and mixtures thereof.

Types of clays considered applicable for best practice in the present invention can also be grouped in general as three-layer types (sheet structures composed of two layers of silica tetrahedrons and one central dodecahedral or trioctahedral) as follows:

- 1. Expanding lattice clays.
  - a. Equidimensional smectite-type clays, including montmorillonite and bentonite
  - b. Elongated smectite clays including saponite.
- Nonexpanding lattice clays including mixed-layer types (ordered stacking of alternate layers of different types);
  - 3. Some clays which by their generic reference fit into more than one of the above-described classification groups.
  - "Generic reference" clays all have the common characteristics of mineral containing aluminum silicate: the major

examples are 1) bentonite, and 2) hectorite

Cation-exchangeable clays can also be synthesized usually from a montmorillonite base clay, so as to possess certain chemical and physical characteristics in order to make such clays valuable in the practice of the present inven-

The above-described clays, especially the bentonlie-type clays, are preferably converted to the sodium form if they are not already in this form. Commercially important representatives of such smedite clays used to make the organoclays of this invention are sodium and calcium bentonte and hectorie. The cation exchange capacity of these smedite-type days can be determined by the well-known mathylene blue method or the armonium acetate method. Smedite-type clays prepared synthetically by either a pneumatolytic or, preferably, a hydrothermal synthesis process may also be used to pregare the novel products of this invention.

The most preferred clays used in the present invention are Wyoming bentonite, particularly sodium bentonite, and hectorie from Hector, California in the Mojave Descr. Bentonite and hectorite clays have high bonding power and reach easily with the specific organic gualernary compounds described hereafter.

In addition, it will be understood that the above listed smeditie-type clays which have been subjected to the application of shear may also be used to make the organoclays of the instant invention. To achieve shearing of the smeditetype clay, the clay is typically dispersed in water at a concentration of from about 0.5 to about 80% by weight. The surry may optionally be first centrifuged to remove non-clay impurities which constitute about 10% to about 95% of time starting clay composition. Of course, if in some cases the clay has previously been freated, such as by the clay vendor, to remove such impurities, the clay as purchased without centrifuging can be formed into a sturry and subjected to shear conditions.

Shear can be imparted to the smecitie-type clay sturry by means of commercially available equipment that is known to impart high shear to the material Illustrative of such equipment are a Manton-Gauin Homogenizer available from the APV Gaulin Company, a Telvmar SD-45 Homogenizer, a Sharples Super Centrifuge available from Sharples Division of Pennwatt Corporation, an other smill available from Gakes Machinery, a Microfluidizer available from Microfluidicer available from Microfluid

The organic cations which are useful in this invention may be selected from a variety of nitrogen-based quaternary materials that are capable of exchanging cations with the selected smettle-type clay. The organic cations which are reacted with smedtle-type clay to prepare the inventive organophilic clays have a positive charge localized on a single nitroen atom within the compound

For this invention, the organic cation is provided by specific quatemary ammonium compounds derived in whole or in part from organic acid - derived esters. This organic cation is provided by a quaternary ammonium compound selected from the group consisting of the following formulae:

wherein  $R_1$  is an alkyl or aralkyl-ester group having 8 to 30 carbon atoms as described below and  $R_2$ ,  $R_3$  and  $R_4$  are independently selected from the group consisting of (i)  $R_1$ , (ii) ong chain linear or branched alkyl (including methyl), allphatic or aromatic groups, kaining 1 to 30 carbon atoms (such groups can also include hydroxylcate groups), (iii) aralkyl groups, such as benzyl and substituted benzyl moieties, including such groups having fused ring moieties having linear chains or branches of 1 to 30 carbon atoms; (iv) any groups such as phenyl and substituted phenyl including fused ring and substituted the properties of the such groups having up to six carbon atoms or hydroxylalkyl groups having up to six carbon atoms or hydroxylalkyl groups having 2 to 6 carbon atoms; and (ivi) hydrogen. M is an anion, typically chloride, mothyl sulfato, acotate, include or bromide.

This quaternary ammonium compound must contain at least one linear or branched, saturated or unsaturated alkyl or aralkyl-ester R, group havino 8 to 30 carbon atoms. Such ester groups are of the general formula:

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where each D is independently selected from the group consisting of H,  $CH_3$  and  $C_2H_5$  and A is an alkyl or aralkyl radical group.

The alkyl and aralkyl-ester groups may be derived from naturally occurring fatty oils including various vegetable including various vegetable and oils, such as como oil, occord, oild, sophean oil, octonated oil, catch or oil and the like, as well as various vegetable and oils or less, the most common of which are occord, oil, sopabean oil and tallow oil (obtained mostly from beef occord) and various vegetable and various vegetable veg

Useful organic ester-derived qualernary ammonium compounds for making the organoclays of this invention include ester qualernary compounds sold by Fina Corporation, hydrogenated tailow ester qualernaries sold by KAO. Chemical Company under the tradename Tetrarium (including Tetrarium) AHT-2 and Tetrariy ACHF 2 and coce ester quaternaries also sold by KAO Corporation under the tradename Quartamin including Quartamin AHT-2 and Quartamin ACHF-2. These quaternaries and similar useful quaternaries made by both these companies, and by other companies, have the general formula.

$$\left[ \begin{array}{c|c} O & H & H \\ \hline & I & & I \\ A^1 - C - O - C - C \\ & & I \\ & H & H \\ \end{array} \right]_b^+ \quad M'$$

where A<sup>1</sup> represents alkyl radicals of either hydrogenated tallow or coconut fatty acids; b and c are 1, 2 or 3 and the total of b and c is always 4: M' is either chloride or methylsulfate.

Particularly preferred for this invention are dimethyl dialkyl-ester quaternary compounds.

Representative examples of useful branched aikly groups include 12-methylsteayl and 12-ethylsteayl. Representative examples of useful branched unsaturated groups include 12-methyloley and 12-ethyloley 18-persentative examples of unbranched saturated groups include launy; stearyl; tridecyl; myristyl; tetradecyl; pertadecyl; hexadecyl and doccsamyl. Representative examples of unbranched, unsaturated and unsubstituted groups include oleyl, linoleyl, linolenyl, soya, coccnut and tallow it is to be noted that natural oils such as soya, cocconut and tallow are in fact mixtures of different carbon chain length elements.

Examples of arallyl, that is benzyl and substituted benzyl moieties, include those materials derived from, e.g., benzyl halides, benzhydryl halides, kirtyl halides, alpha-halo-alpha-phenylalkanes wherein the alkyl chan has trom 1 to 30 carbon atoms, such as 1-halo-1-phenylethane, 1-halo-1-phenylpropane, and 1-halo-1-phenyloctadezane, substituted benzyl moieties, such as those derived from ortho-, meta- and para-chlorobenzyl halides, para-methocybenzyl halides, and cribo-, meta- and para-chlorobenzyl halides, para-methocybenzyl halides, and ortho-, meta- and para-chlorobenzyl halides wherein the alkyl chain contains from 1 to 30 carbon atoms, and fused ring benzyl-type moieties, such as those derived from 2-halomethylnapthietiene, 9-halomethylnathracene and 9-halomethylphenathrace, wherein the halogenated group comprises chloro-, bromo-, lodo-, or any other such group which serves as a leading group in the nucleophilic attack of the benzyl-type moiety so that the nucleophilic replaces the leading group on the benzyl-type moiety.

Examples of any igroups that are useful include phenryl and substituted phenryl. Nathyl and N.N-dialkyl anilines, wherein the skily groups contain between 1 and 30 carbon atoms, orthor, mate- and para-aikyl phenryl, wherein the skily group contains between 1 and 30 carbon atoms, 2, 3-, and 4-halophenryl wherein the halo group is defined as chitor-, bromo-, or iodo-, and 2, 3-, and 4-carboxyphenryl and esters thereof, where the alcohol of the ester is derived from an alkyl atochol, wherein the alkyl group contains between 1 and 30 carbon atoms, any such as a phenryl carafixly such as benryl alcohols, fused ring any limiteties such as naphthleane.

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anthracene, and phenanthrene

Beta, gamma unsaturated alkyl groups which may be included in organic catlon component of the organophilic clay gellants of the invention may be selected from a wide range of maletials well known in the ari. These compounds may be cyclic or acyclic unsubstituted or substituted with alighatic radicals containing up 50 carbon aloms such that the total number of alighatic carbons on the beta, gamma unsaturated radical is 6 or less. The beta, gamma unsaturated alkyl radical may be substituted with an aromatic ring that likewise is conjugated with the unsaturated beta, gamma molely of the beta, gamma radical may be substituted with on a signatic radicals and aromatic rings.

A representative examples of beta, gamma unsaturated alkyl groups include 2-cyclchexenyl. Representative examples of acyclic beta, gamma unsaturated alkyl groups containing 6 or less carbon atoms include propargyl, alkyl (2-propenyl, conyl (2-butenyl), 2-pentenyl; 2-pentenyl; 2-pentenyl; 3-methyl-2-butenyl; 3-methyl-2-pentenyl; 2-pentenyl; 2-pentenyl;

Hydroxyalikyl groups may be selected from a hydroxyl substituted alliphatic radical wherein the hydroxyl is not substituted at the carbon atom adjacent to the positively charged atom the group has thorne 2 to 8 alliphatic carbon atoms. The alkyl group may be substituted with an aromatic ring independently from the 2 to 6 alliphatic carbons. The selection of the

The groups found to be most effective in producing the organic ester-derived quaternary ammonium compounds of the type found useful in this invention, in addition to one or more B<sub>1</sub> organic acid - derived ester groups of the type described are, (a) one or more methyl groups, (c) one or more benzyl groups, (c) one or more long-thoran lakyl groups, and combinations thereof. Most preferred B<sub>1</sub> groups are those derived from occonut oil. Specific quaternaries found most useful are docco- esters of dimethyl diethenol ammonium chloride

A broad variety of quaternary ammonium compounds are disclosed in U.S. Patent No. 4,141,841 issued to Proctor & Gamble Company - see also U.S. Patent No. 3,862,058 also issued to Proctor & Gamble

The preparation of quaternary compounds used to make organophilic clays of this invention can be achieved by techniques well-known in the art. For example, when preparing a quaternary ammonium compound of the type described one skilled in the art may begin with a readily available teritary alkanolamine, such as triethenolamine, methyl or diethyl diethanolamine, diethyl for diethyl proparolamine, which may be reacted with one or more falty acids under esterification reaction conditions, the practitioner will recognize that the molar ratio of fatty acid that react with the hydroxyl moleties of the alkanol amine preferably should be at most 11, and can be less than 1:1. When the ratio is less than 1:1, then the resulting seits groups will be statistically distributed among the alkanol groups. If, on the other hand, the ratio of latty acid to alkanol molettes is greater than 1:1, some amount of the fatty acid will remain unesterfied following the esterification reaction, which amount may or may not have some small adverse effect on subsequent reactions of the product of this reaction.

Following the esterification of some of all of the hydroxyl moieties of the alkanol groups of the tertiary amine, the reaction product may, optionally, be reduced with a dilluent, typically water and/or a low molecular weight alcohol such as methanol, ethanol, 2-propanol or butanol, and reacted with an alkylating agent under conditions well known to one skilled in the art to form the quaternary ammonium compounds useful in making the organoclays of this invention.

The organic anion(s) optionally employed in the products of the invention may be selected from a wide range of materials that are capable of reacting with the organic cations in order to form an organic cation/organic anion complex. The molecular weight of the organic anion is preferably 3,000 or less, and more preferably 1,000 or less, and contains at least one anionic molety per molecule so as to permit the formation of the organic cation/organic anion complex which then becomes intercalated between the cite y platelets

Preferred organic anions are derived from carboxylic acids, such as stearic acid, oleic acid, palmitic acid, succinic acid, tartaric acid, etc.; sulfonic acids; and alkyl sulfates, such as the lauryl half ester of sulfuric acid and mixtures thereof.

The organic anion, which may include mixtures of organic anions, is reacted with the organic cation and smectitetype clay to form the desired organophilic clay gollant. The organic anion may be added to the reaction mixture in acid or saft form. Exemplary of the latter form are alkali metal safts, alkaline earth safts, ammonium and organic amines.

Representative salts of the organic anion are those formed with hydrogen, lithium, sodium, potassium, magnesium, calcium, barium, ammonium and organic amines such as ethanolamine, diethanolamine, triethanolamine, methyldiethanolamine, diethylamine, and so forth, and mixtures thereof. The most preferred salt form is with sodium.

The amount of organic anion optionally reacted with the smectite-type clay and the organic cation must be sufficient to the amount of organic actions to organic anion in the range of from about 1.70.1.0 to about 51.0, preferably from about 3.0.1.0 to about 15.1.0. The most prefered ranges depend on the particular organic cations and optional organic anion utilized and the intended environment of use and can be determined by experimentation guided by the information set forth above. Illustrative perfents which describe suitable organic anions that may be corrected with the organic cations and the smectite type clay in order to form the organic pricipal clay include commonly assigned IIS. Patent Nac. 4.12.0.18 and 4.34.0.075.

The present invention also contemplates a process for preparing an organophilic clay gellant using the specified quaternary compounds

The organophilic clays of this invention may be prepared by admixing the clay organic cation, optional organic anion and water rogether, preferably at amperatures within the range from 20° to 100°C, and most preferably from 35° to 50°C for a period of time sufficient for the organic compounds to react with the clay. The reaction is followed by filtering, washing, drying and grinding. The organic cation may be added simultaneously or at separate intervals in any order.

The clay is preferably dispersed in water at a concentration from about 1 to 80%, most preferably, from 2 to 8% Optionally, the sturry may be centrifuged to remove non-clay impurities which may constitute about 10 to 50% of the starting clay composition.

The amount of the quaternary ammonium compound or compounds added to the smecifie clay for purposes of this invention must be sufficient to impart to the clay the improved gelling and dispersion characteristics. This amount is defined as the milliequivalent ratio, which is the amount of milliequivalents (m.e.) per 100 grams of natural clay without impurities. Such ratio and its calculation are well known in the art.

The organophilic clay gellants prepared according to the invention may be used as meobigical additives in nonaqueous compositions such as paints varniehes, enamels, waxes, paint-varnish lacquer remover, oil base drilling fluids, lubricating grease, inks, polyyester reains, epoxy resins, mastices, adhesives, sealants, cosmetics, detergents, and the like. The organocialy can be added to these systems by commonly known matter including medium speed dispersers, colloid mills, roller mills, and ball mills.

While not wishing to be bound by any theoretical mechanism, it is believed that organoclays of the instant invention will be biodegradable - or more properly, environmentally degraded - as a result of hypothysis of the ester linkage of the quaternaries of the instant invention. It is well known that the reaction(s) of a quaternary ammonium choinde with a smeetite-type clay to form an organoclay is an equilibrium reaction, and that, for the reaction to proceed to completion (e. quantitative consumption of the quaternary ammonium choinde), at least one of the moistless statched to the quaternary ammonium cations of shorter chain lengths are in equilibrium with the raw materials for the organoclay rise believed that this dependence on alkyl chain length is due to van der Waas forces between the alkyl chain and the surface of the clay platelet; only when the van der Waasl forces of attraction are greater than the solution energy of the quaternary is the quaternary quantitatively consumed in the reaccion(s) to form the organoclay.

The fatty acid(e) seter(s) of the quaternary variance was considered to the instant invention provide the required chain from the fatty acid(e) seter(s) of the quaternary ammonium cations of the instant invention provide the required chain length to form stable organociays. However, once the seter inkage(s) hydrolyse, the remaining quaternary cations of not possess the requisite long chain alkyl mioley, as a result, said hydrolyse product (i.e., the organociay with a hydrolyzed ester quaternary cation) will equilibrate with its environment in which cations other than the hydrolyzed ester quaternary cation are likely to be more prevalent returning the organociacy, ultimately, to a totally inorganic form. The quaternary ammonium cation released from the hydrolyzed ester quaternary organoclay may further degrade by other mechanicisms.

The organoclays of this invention can be used in combination with other materials including organoclays made with conventional prior and quaternary compounds. Consequently, the invention also provides non-aqueous organic and solvent compositions thickened with the above-indicated organophilic clay gellant. Thus, a third aspect of the invention relates to a non-acueous (fluid system which comprises.)

(a) a non-aqueous liquid composition such as paint, coatings, drilling fluids, ink or similar materials; and
 (b) an organophilic clay gellant comprising the reaction product of:

- (i) a smectite-type clay having a cation exchange capacity of at least about 75 milliequivalents per 100 grams of clay, 100% active clay basis;
- (ii) an organic cation or cations, as described, in an amount sufficient to satisfy at least about 75%, preferably about 100%, of the cation exchange capacity of the smectite-type clay and of any optional organic anion(s), and optionally optional organic anion(s).
- (iii) one or more organic anion(s) that are capable of reacting with the organic cation or cations to form a complex as described previously.

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The organophilic city complexes of the invention are added to the non-aqueous compositions in amounts sufficient to obtain the desired redeological proparties. Amounts of the organophilic city complexes in the non-aqueous compositions are from about 0.01% to 15%, preferably from about 0.3% to 5%, based on the total weight of the non-aqueous filled section.

The following examples are given to illustrate the invention, but are not deemed to be limiting thereof. All percentages given throughout the specification are based upon weight unless otherwise indicated.

## EXAMPLE 1

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This example and Examples 2 to 6 illustrates the preparation of organoclays according to the present invention. In this example, 45.00 grams of beneficiated un-sheared Wyoming behonite lotely sutury was diluted with water to form a 2% by weight dilute slurry of benfonte in water. The slurry was heated to 65°C in a reaction flask equipped with a stirrer, thermometer and addition funnel 28.5 4 grams of al diococo eater of dimethy diethanol ammonium methysulphate quaternary compound, commercially available as TETRANYL ACHF 2 from Kao Corporation, was added to the clay slurry. The mixture was sitred for 50 minutes at 65°C. The product was filtered through a Buchner funnel to collect the solids. The wat solids were resiturized in 1500 grams of water at 65°C for 20 minutes and then recollected on a Buchner furnel. The organocisty "filteracket" was drund at 45°C to 15 hours.

#### **EXAMPLE 2**

The procedure of Example 1 was repeated, except that 34.62 grams dihydrogenated tallow ester of dimethyl diethanol ammonium methylsulphate, commercially available as TETRANYL AHT-2 from Kao Corporation was used.

## COMPARATIVE EXAMPLE A

For comparative purposes, the procedure of Example 1 was repeated, except that 24.15 grams of dimethyl dihydrogenated tallow armonium chloride, a quaternary compound commonly used to make a variety of commercial orq

# 30 EXAMPLES 3-6

The compositions were prepared according to the procedure of Example 1, except that 28.54 grams of the following quaternary ammonium compounds were used:

Example 3 - dihydrogenated tallow ester of dimethyl diethanol ammonium chloride

Example 4 - dihydrogenated tallow ester of methyl triethanol ammonium methylsulphate

Example 5 - dicoco ester of methyl triethanol ammonium methylsulphate Example 6 - dicoco ester of dimethyl diethanol ammonium chloride

Example of Global district of differing decidation and months in entone

## 40 EXAMPLES 7-19

A typical organic system paint formulation was prepared according to Formulation 1.

#### Formulation 1

Long Oil Alkyd Paint

| Ingredients   | Description  | Manufacturer | Amount (Pounds)                           |
|---|--|--------------|---|
| Millbase  |  |              |   |
| Beckosol 10-060<br>Mineral Spirits 66/3<br>Organoclay<br>MeOH/H2O 95/5<br>KRONOS 2101 | Long oil alkyd<br>Solvent<br>Polar activator<br>Titanium Dioxide | Reichhold    | 105.76<br>70.60<br>7.17<br>2.39<br>325.00 |
|   |  |              |   |

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#### (continued)

| Ingredients          | Description     | Manufacturer | Amount (Pounds) |
|----------------------|-----------------|--------------|-----------------|
| Letdown              |                 |              |                 |
| Beckosol 10-060      | Long oil alkyd  | Reichhold    | 445.90          |
| 6% ZR Nuxtra         | Drier           | Hüls         | 10,30           |
| 6% Co Nuxtra         | Drier           | Hüls         | 3.42            |
| EXKIN #2             | Anti skin agent | Hüls         | 2.00            |
| Mineral spirits 66/3 | Solvent         | 1            | 54.70           |
| • ***                |                 | Į.           | 1027.24         |

Each of the organoclay samples prepared in Examples 1 to 6, Comparative Example A and Bentone 34, were dispersed into Formulation 1 at a loading of 0.7% using a Dispersmat disperser B34 (Bentone 34) is a commercial organoclay made by Rheox Inc. using a nitrile quaternary derived from tallow. The results demonstrate the viscosity and other typical coating properties provided when the novel organoclays of the invention when used in an alkyd resin paint formulation. The various tests employed were standard laboratory measurements conventionally used in determining the effectiveness of organoclays in providing rheological properties to organic liquids.

# Paint Properties:

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| 25 | Example<br>(Composition)                        | FOG        | Stormer Viscosity<br>(KU) | Brookfield 10<br>RPM (cP) | Viscosity 100<br>RPM (cP) | T.I. | Sag<br>(Mil) | Gloss    |
|----|---|------------|---------------------------|---------------------------|---------------------------|------|--------------|----------|
|    | 7 (Example 1)                                   | 7BA        | 91                        | 2120                      | 1264                      | 1.68 | 4.8          | 90       |
|    | 8 (2)   | 7BA        | 86                        | 1680                      | 1040                      | 1.62 | 4.2          | 91       |
|    | 9 (Example 3                                    | 7BA        | 86                        | 1680                      | 1088                      | 1.54 | 3.9          | 94       |
| 30 | Comp A)<br>10 (B34)<br>11 (no additive<br>used) | 7BA<br>7BA | 89<br>82                  | 1800<br>800               | 1144<br>800               | 1.57 | 5.0<br>3.0   | 89<br>91 |

| 35 |                          |     |                           |                           |                           |      |              |       |
|----|--------------------------|-----|---------------------------|---------------------------|---------------------------|------|--------------|-------|
|    | Example<br>(Composition) | FOG | Stormer Viscosity<br>(KU) | Brookfield 10<br>RPM (cP) | Viscosity 100<br>RPM (cP) | T.I. | Sag<br>(Mil) | Gloss |
|    | 12 (4)                   | 6C  | . 94                      | 2100                      | 1384                      | 1,52 | 5            | 90    |
|    | 13 (5)                   | 6BC | 89                        | 1700                      | 1160                      | 1.47 | 5            | 88    |
| 40 | 14 (B34)                 | 7AB | 94                        | 2220                      | 1414                      | 1.57 | 5            | 89    |
|    | 15 (no additive          | 7AB | 82                        | 800                       | 800                       | 1.00 | 3            | 91    |
|    | used)                    |     | 1                         | 1                         |                           | 1    | ĺ            |       |

| 45 |                          |     |                           |                           |                           |      |              |       |
|----|--------------------------|-----|---------------------------|---------------------------|---------------------------|------|--------------|-------|
|    | Example<br>(Composition) | FOG | Stormer Viscosity<br>(KU) | Brookfield 10<br>RPM (cP) | Viscosity 100<br>RPM (cP) | T.I. | Sag<br>(Mil) | Gloss |
|    | 16 (3)                   | 7B  | 91                        | 2900                      | 1430                      | 2.03 | 5.6          |       |
|    | 17 (6)                   | 7B  | 86                        | 2000                      | 1150                      | 1.74 | 4.8          |       |
| 50 | 18 (B34)                 | 7B  | 87                        | 2080                      | 1140                      | 1.82 | 4.9          |       |
|    | 19 (Blank)               | 7B  | 75                        | 700                       | 620                       | 1.13 | 3.0          |       |

Discussion of Results: Many of the above results demonstrate that organoclays made according to this invention are superior to conventional organoclays - compare examples 7 to 9 and 16 to 18. While some of the examples shown equal or slightly diminished effectiveness, all inventive examples are technically and commercially acceptable Based on the foregoing results, it is apparent that the organophilic clays provided by the invention are highly

effective in improving the rheological properties of non-aqueous systems.

The invention thus being described, it will be obvious that the same may be varied in many ways. Such variations are not to be regarded as a departure from the spirit and scope of the invention and all such modifications are intended to be included within the scope of the claims.

#### Claims

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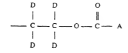
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- An organophilic clay for providing rheological properties to non-aqueous fluid systems comprising the reaction product of:
  - (a) one or more smectite-type clays having a cation exchange capacity of at least about 75 milliequivalents per 100 grams of clay, 100% clay, active clay basis; and
  - (b) one or more quaternary ammonium compounds derived from an organic acid-derived ester in an amount sufficient to satisfy at least about 75% of the cation exchange capacity of the smectite-type clay.
- 2. The organophilic clay of claim 1, wherein one or more organic anions are included in the reaction product to form a complex, the quaternary armmonium compound(s) being present in amount sufficient to satisfy at least about 75% of the cation exchange capacity of the smectite-type clay and of the organic anion, and said organic anion (s) preferably being derived from carboxytic acids, sulfonic acids, silyl sultates or mixtures thereof
- The organophilic clay of claim 1 or 2, wherein said smectite-type clay is selected from bentonite, hectorite and mixtures thereof.
- 4. The organophilic clay of any of claims 1 to 3, wherein one or more of the quaternary ammonium compounds have the formula:

$$\begin{array}{c|c}
R_1 & M^- \\
R_2 - N - R_4 \\
R_3
\end{array}$$

wherein R<sub>1</sub> is an alkyl or aralkyl-seter group having 8 to 30 carbon atoms and R<sub>2</sub>. R<sub>3</sub> and R<sub>4</sub>, are each independently selected from (9 R<sub>4</sub>, (ii) long chain lineaer or branched alkyl (notuding methlyk), alphalic or aromatic groups having 1 to 30 carbon atoms optionally substituted by hydroxylated groups; (iii) aralkyl groups such as benzyl and substituted brary moisties, including such groups having fused ring moisties having linear chains or branches of 1 to 30 carbon atoms; (iv) anyl groups such as phenyl and substituted pharply including fused ring aromatic austituted the control of the con

5. The organophilic clay of claim 4 wherein the or each R<sub>1</sub> group has the formula:



wherein each D is independently selected from H,  $CH_3$  and  $C_2H_5$  and A is an alkyl or aralkyl radical group which group has been derived from naturally occurring oils selected from occount oil, soybean oil and tallow

6. The organophilic clay of any of claims 1 to 3 wherein one or more of the quaternary ammonium compounds have

- where A<sup>1</sup> represents an alkyl radical derived from hydrogenated tallow or coconut fatty acids ib and c are each 1, 2 or 3 and the total of b and c is always 4; and M- is either chloride or methyl sulfate.
  - 7. The organophilic clay of any of claims 1 to 6 wherein the quaternary ammonium compound is selected from a dicoco ester of dimethyl diethanol ammonium methylsulphate, a dicoco ester of methyl tiethanol ammonium methylsulphate a dicoco ester of dimethyl diethanol ammonium chloride, a dhydrogenated tallow ester of dimethyl diethanol ammonium methylsulphate and a dhydrogenated tallow seter of dimethyl diethanol ammonium methylsulphate and a dhydrogenated tallow seter of dimethyl diethanol ammonium methylsulphate.
  - 8. A process for preparing an organophilic clay according to any of claims 1 to 7, which process comprises:
    - (a) preparing an aqueous slurry of the smectite-type clay having a cation exchange capacity of at least about 75 milliequivalents per 100 grams of 100% clay, active clay basis;
    - (b) heating said slurry to a temperature of from about 20°C to 100°C
    - (c) adding to said sturry the quatermary ammonium compound(s) and optionally one or more organic anions as defined in claim 2, the quatermary ammonium compound(s) being present in an amount sufficient to satisfy at least about 75% of the cation exchange capacity of the smedite-type clay and of the organic anion if present.
    - (d) reacting the resulting mixture for a sufficient time to form an organophilic clay; and
    - (e) recovering the organophilic clay.
- 95 9. A non-aqueous fluid system coniprising a non-aqueous composition and an organophilic clay according to any of claims 1 to 7.
  - 10. The non-aqueous fluid system of claim 9, wherein said non-aqueous composition is selected from paints, coatings, varnishes, enamels, waxes, paint-varnish, lacquer remover oil base drilling fluids, greases, inks, polyester resins, epoxy resins mastices, adhesives, sealants, cosmetics and detergents.
  - 11. The non-aqueous fluid system of claim 9 or 10, wherein said organophilic clay is present in an amount of from about 0.01% to 15% based on the total weight of said non-aqueous fluid system.

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European Pai Office

#### EUROPEAN SEARCH REPORT

Application Number EP 97 65 0004

| Category                                    | Citation of document with<br>of relevant  | indication, where appropriate,                              | Relevant<br>to classo  | CLASSIFICATION OF THE<br>APPLICATION (Inc.Cl.6)  |  |  |  |
|---|---|---|--|--|--|--|--|
| x<br>x<br>x                                 | EP 0 204 240 A (HC<br>* claims 1,3,4,6 *<br>* page 5, line 31<br>* page 6, line 24  | ECHST AG) - page 6, line 7 * - line 28; claim 8 *           | 1,3<br>8<br>9-11   | C01B33/44<br>C09D5/04<br>C09D9/00<br>C09K7/02  |  |  |  |
| x<br>x<br>x                                 | DE 34 34 983 A (HO<br>claim 1 *<br>* page 6, line 5 -<br>* page 7, line 1 -<br>* page 8, line 11<br>* page 9, line 11 -                               | line 7 * line 4 * - line 20 *                               | 8<br>9-11  | C10M113/16<br>C09D11/06<br>C08K3/34<br>C09D5/34<br>C09J11/04<br>C09K3/10<br>A61K7/00<br>C11D3/12 |  |  |  |
| A   | DE 38 06 548 A (KA<br>CHUO KENKYUSHO)<br>* page 9, line 33  | BUSHIKI KAISHA TOYOTA<br>- line 39 *                        | 1  |  |  |  |  |
| A   | LU 83 759 A (NL IN<br>* claims 1,9 *<br>* page 11, line 12  | DUSTRIES, INC.) - page 12, line 16                          | 2  |  |  |  |  |
| A, A  | * claim 12 *<br>& US 4 434 075 A  |   | 8  | TECHNICAL PIELDS<br>SEARCHED (Int.Cl.6)  |  |  |  |
| A   | GB 2 090 584 A (NL * claims 1,7,10 * * page 3, line 22 * claims 11,19,20 & US 4 412 018 A   | - line 50 *   | 2  | C01B   |  |  |  |
| , A   | EP 0 542 266 A (RH<br>& US 5 336 647 A  | EOX INTERNATIONAL, IN                                       | ic.)   |  |  |  |  |
| A, A  | GB 2 017 072 A (NL<br>& US 4 208 218 A  | INDUSTRIES, INC.)   |  |  |  |  |  |
|   | The present search report has   | ocen drawn up for all claims  Date of completes of the care |  |  |  |  |  |
|   | THE HAGUE   |   | oion. J  |  |  |  |  |
| λ : parti<br>) · parti<br>desu<br>4 · terni | ATT GORY OF CITED DOCUME<br>marts relevant if taken alone<br>meants relevant if combined with an<br>ment of the same category<br>sological background | E : cattler pat after the fi  D : document L : document     | erinciple underlying the<br>ent document, but public<br>ling date<br>cited in the application<br>cited for other reasons | invention<br>ined so, or   |  |  |  |
|   | entre dischium<br>mediar forument   | 4 : member of   | the same patent family   | corresponding  |  |  |  |

